

The ontogeny of continuous quantity discrimination in zebrafish larvae (*Danio rerio*)

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22 **ABSTRACT**

23 Several studies have investigated the ontogeny of the capacity to discriminate between discrete
24 numerical information in human and non-human animals. Contrarily, less attention has been
25 devoted to the development of the capacity to discriminate continuous quantities. Recently, we set
26 up a fast procedure for screening continuous quantity abilities in adult individuals of an animal
27 model in neurodevelopmental research, the zebrafish. Two different sized holes are presented in a
28 wall that divides the home tank in two halves and the spontaneous preference of fish for passing
29 through the larger hole is exploited to measure their discrimination ability. We tested zebrafish
30 larvae in the first, second and third week of life varying the relative size of the smaller circle (0.60,
31 0.75, 0.86, 0.91 area ratio). We found that the number of passages increased across the age. The
32 capacity to discriminate the larger hole decreased as the ratio between the areas increased. No
33 difference in accuracy was found as a function of age. The accuracy of larval zebrafish almost
34 overlaps that found in adults in a previous study, suggesting a limited role of maturation and
35 experience on the ability to estimate areas in this species.

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38 **KEYWORDS:** Size discrimination; Zebrafish; Continuous extent; Spatial abilities.

39

40 INTRODUCTION

41 Cultural (Butterworth et al. 2011; Pica et al. 2004), cognitive (Price et al. 2012; Revkin et al.
42 2008) and developmental (Izard et al. 2009; Xu and Spelke 2000) psychology have provided
43 multiple evidence for the existence of quantificational abilities that do not rely on language and
44 culture. Such cognitive abilities, commonly referred to ‘non-symbolic’ quantificational skills,
45 permit us to quickly solve many problems in everyday life, such as avoiding longer queues,
46 selecting the shorter distance or the larger amount of food. Given the multiple advantages in terms
47 of fitness and survival in the natural environment, quantification abilities are also shared by many
48 vertebrate species (reviewed in Henik 2016).

49 There are two broad categories of quantificational skills, namely, estimation of discrete
50 quantities (i.e. numerosity), and estimation of continuous quantities (e.g. distance, length, area,
51 weight or duration). For what concerns the capacity to estimate numerosity, there is evidence that in
52 our species it is innate and that it increases in precision across development even before the
53 emergence of language, which occurs after 12-13 months of age (Fenson et al. 1994). Newborns can
54 discriminate between two numbers up to a 0.33 numerical ratio (4 vs. 12 objects) (Izard et al. 2009);
55 6-months-old infants can discriminate a 0.50 ratio (8 vs. 16 objects) but not a 0.67 (8 vs. 12 objects)
56 (Xu and Spelke 2000). Such ratio can be discriminated only later, at 10 months of age (Xu and
57 Arriaga 2007). Our numerical acuity keeps improving throughout childhood, with 6-years olds
58 discriminating a 0.83 ratio and adults discriminating a 0.90 ratio (Halberda et al. 2008; see Cordes
59 and Brannon 2008 for a review).

60 Numerical abilities have been investigated in more than 50 species representing a wide
61 range of vertebrate and invertebrate groups (e.g. Agrillo and Bisazza 2010; Beran and Parrish
62 2016). Developmental research is however limited to a few species. Inborn numerical abilities were
63 found in domestic chicks tested with multiple experimental paradigms (e.g. Rugani et al. 2007,
64 2009, 2015). A small teleost fish, the guppy (*Poecilia reticulata*), demonstrated an innate ability to
65 discriminate small numbers of social companions while their capacity to discriminate larger

66 numerosities emerges later, as a result of both maturation and experience (Bisazza et al. 2010).
67 Another study, using an operant conditioning procedure, showed that newborn guppies can
68 discriminate between large numerosities in the first days of life (Piffer et al. 2013) but only if they
69 could use multiple cues, such as number and cumulative surface area (Miletto Petrazzini et al.
70 2014).

71 For what concern continuous quantities, there is evidence that in humans this capacity
72 improves across development too (reviewed in Feigenson 2007): 6-month old infants tested for their
73 ability to discriminate between two areas could discriminate a 0.50 but not a 0.67 size ratio
74 (Brannon et al. 2006). The latter ratio is discriminated at 10 months of age (Feigenson et al. 2002).
75 Studies of continuous quantity estimation are rare in other species. Most evidences are indirect; for
76 instance, it was found that different fish species discriminate their potential mates depending on
77 their size (e.g. Bisazza et al. 2000; Dosen and Montgomerie 2004); also, size-assortative social
78 grouping is widespread across animal species and seems the consequence of the ability to gauge
79 conspecifics size (e.g. Barry et al. 2014; Pitcher 1986). Continuous quantity estimation abilities
80 have been directly investigated in a half dozen species including sea lions (*Zalophus californianus*),
81 carrion crows (*Corvus corone*), ruin lizards (*Podarcis sicula*) and Hermann's tortoises (*Testudo*
82 *hermanni*) (Gazzola et al. 2018; Miletto Petrazzini et al 2017; Moll and Nieder 2014; Schusterman
83 et al. 1965). As concerns fish, a recent work investigated in the guppy the ability to discriminate
84 between different numbers of food items or between food items that differed in size. The study
85 demonstrated that the guppies were able to discriminate both discrete and continuous quantities but
86 were much more accurate in the latter task (Lucon-Xiccato et al. 2015). Although this ability is
87 likely to have several functions even in young animals (e.g. social grouping, food choice, fighting
88 assessment), there is a complete lack of studies regarding the ontogeny and the development of
89 continuous quantity abilities in non-human species.

90 In humans, neuro-developmental and neuro-degenerative disorders are associated with a
91 decline in numerical abilities and in the ability to estimate continuous quantities, for example the

size, the weight and the distance of an object or the duration of an event (Barabassy et al. 2010; Brand et al. 2003; Girelli et al. 1999).

The zebrafish (*Danio rerio*) represents a well-established model in developmental, genetic and pharmacological research. This species is rapidly becoming a model for human neurodevelopmental and neurodegenerative disorders (e.g. Brennan, 2011; Kalueff et al. 2014; Xi et al. 2011). The quasi-totality of these studies is conducted on embryos or in larvae in the first weeks of life. Larvae can be readily obtained in large numbers, allowing large-scale screening of neuroactive compounds and genotypes (e.g. Norton, 2013; Richendrfer et al. 2012). The rapid development of zebrafish favours the investigation of the developmental mechanisms whereas the transparency of the larvae allows the study of the neural circuits underlying behavioural and cognitive functions through in vivo calcium imaging (e.g. Sarvaiya et al. 2014). For these reasons, tools for modelling brain diseases in larval zebrafish are becoming increasingly available (e.g. Buckley et al. 2008; Paquet et al. 2009). Although cognitive deficits are important markers for many human neuropathologies (e.g. Levin et al 1989; Morris et al. 2001), there is currently a complete lack of protocols for measuring quantity estimation abilities in early developmental stages of this species.

Recently, we developed a simple procedure to test continuous quantity discrimination in adult zebrafish. The procedure is based on a spontaneous preference of fish to pass through the larger available breach (Santacà et al. 2020). We recorded small groups of zebrafish, composed of six adult females, that could move between the two compartments of their housing tank passing through a divider provided with two holes differing in size. In a first experiment, zebrafish significantly discriminated area ratios from 0.60 to 0.91 but their discrimination performance was found to decrease as the relative size of the smaller hole increased, in agreement with Weber's law. In a second experiment, we measured the performance of individually tested fish. On average, performance of individually tested fish largely overlapped performance of fish tested in clusters. However, a large inter-individual variation in performance was observed in this experiment.

117 Very little is known about the quantificational abilities of zebrafish in the early stage of life.
118 Two studies have found that zebrafish larvae were attracted by a small moving dot (a potential prey)
119 but avoided a very large moving dot (a potential predator) (Bianco et al. 2011; Barker and Baier
120 2015) meaning that they are provided with at least some rudimental size discrimination ability.
121 Neither of the two studies, however, investigated the developmental trajectory of such skill, nor
122 they attempted to assess the threshold of discrimination.

123 Our study aimed to fill the gap in knowledge regarding the ontogeny and the developmental
124 trajectory of continuous quantity discrimination, adapting the procedure developed for adults to
125 study larval zebrafish. We observed the ability of larval fish to discriminate the larger hole to move
126 from one compartment to another of their home tank. Zebrafish larvae were tested at three different
127 ages, in the first, in the second and in the third week of life (7-, 14- and 21-days post fertilization
128 respectively) varying the ratio of the smaller to the larger hole (0.60, 0.75, 0.86 and 0.91 area
129 ratios). These information could also have practical applications as they may allow to study the
130 cognitive deficits associated with neuropathologies in larval stages instead of adult zebrafish.

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132

133 **METHODS**

134 *Subjects*

135 We tested one hundred and eighty zebrafish larvae (*Danio rerio*), in particular 30 groups of
136 6 larvae each for three different ages. Throughout this manuscript we used the standard age
137 classification for zebrafish studies which start with the fertilization day and is expressed in days
138 post fertilization or dpf (e.g. Bilotta and Saszik 2001). Since zebrafish hatch approximately 72
139 hours after fertilization, age from birth can be calculated subtracting three days from the number of
140 dpf. We tested 10 groups larvae at 7 dpf, 10 groups of 14 dpf, and 21 dpf. The sex of the larvae was
141 undetermined as sexual differentiation completes much later around week 11-12 post fertilization
142 (Maac et al. 2003). The larvae used in our experiment originated from many different breedings
143 from a wildtype strain bought by a local supplier in 2018 and maintained in our laboratory in a large
144 population (>200 fish). The larvae were raised in several petri dishes in a solution of Fish Water 1x
145 (0.5 mM NaH₂PO₄*H₂O, 0.5 mM Na₂HPO₄*H₂O, 1.5 gr Instant Ocean, 1 L de-ionized H₂O) and
146 Methylene blue (0.0016gr/l). Until the beginning of the experiment, they were housed at a density
147 of approximately 30 individuals each petri dish in the same room maintained at a temperature of
148 28.5 ± 1 °C. The room was illuminated according to a 14:10 h light:dark cycle. Larvae were fed
149 twice a day with dry food (particle size: 0.75 mm) from the age of 6 dpf.

150 To avoid stressing the subjects, the standard length (from the snout to the caudal peduncle)
151 and the width (from the left to the right side of the head) of each larvae was obtained from video
152 recording of the experiment using the computer software Image J (<https://imagej.nih.gov/ij/>). For
153 each larvae, we calculated the mean from three different video frames. The 7 dpf larvae were on
154 average 3.962 ± 0.092 mm (mean ± SD) long and 0.612 ± 0.048 mm wide, the 14 dpf larvae were
155 4.397 ± 0.118 mm long and 0.732 ± 0.088 mm wide whereas the 21 dpf larvae were 4.565 ± 0.124
156 mm long and 0.918 ± 0.073 mm wide. The height at nape of the larvae could not be obtained from
157 the video recordings; therefore, it was measured from a sample of larvae (n = 30) that did not
158 participate in the study and consisted in the distance from ventral to dorsal, immediately posterior of

159 head, perpendicular to the axis defined by the standard length. The 7 dpf larvae were on average
160 0.731 ± 0.061 mm (mean \pm SD), the 14 dpf larvae were 0.914 ± 0.098 mm whereas the 21 dpf
161 larvae were 1.022 ± 0.051 mm high.

162 *Apparatus*

163 The experiment was conducted in an hourglass-shaped apparatus (12 x 4.8 cm and 4 cm
164 high) filled with 3.5 cm of Fish Water 1x (Figure 1). The apparatus was 3D printed with white PLA
165 material. A central corridor (4.3 cm in length) connected the frontal and the posterior
166 compartments. In the middle of the corridor, larvae could perform the task that consisted in the
167 spontaneous passing through one of two holes of a plastic panel (3 x 3.2 cm; Figure 1). The panels
168 were 3D printed with grey PLA material. Larvae could move from one compartment to the other
169 one of the apparatus passing through the holes. One 0.72 W LED lamp was placed 1 cm above the
170 frontal compartment, and one above the posterior one. We used four identical apparatuses at the
171 same time. One video camera was placed above the central corridors of two adjacent apparatuses.

172 *Procedure and stimuli*

173 Since zebrafish is a highly social species, social isolation could alter the development of
174 their normal behavioural repertoire, we studied fish in small social groups, the same condition in
175 which this species is expected to solve this type of problem in nature (Forsatkar et al. 2017). The
176 procedure consisted of two phases: a habituation phase and a test phase; both phases lasted two
177 days. On day 1 of the habituation phase, six same-age larvae (7 dpf, 14 dpf or 21 dpf) were
178 randomly selected from different maintenance petri dishes and were inserted in an experimental
179 apparatus. On day 2, a grey panel with one central big hole (0.7 cm in diameter) was inserted in the
180 middle of the apparatus to make the fish familiar with the procedure (i.e. passing through a hole to
181 move from one side to the other). During these days, fish could habituate to the tank and swim
182 through the hole. We video recorded the second day to ensure that the larvae had habituated to the
183 experimental procedure passing through the hole. On the third and fourth day (test phase), four
184 different panels with four types of size discrimination trials were randomly presented: ratio 0.60,

ratio 0.75, ratio 0.86 and ratio 0.91 between the areas of the two holes (Figure 2). The value of the ratio represents the relative size of the smaller hole compared to the larger one that was maintained fixed in all four discriminations. A pilot experiment indicated that larvae easily pass through holes of 0.4 cm in diameter. However, since larvae have a poorer visual acuity than the adults (see discussion), to favour simultaneous visibility of the two holes even from a distance, we used holes above 0.6 cm of diameter (see Table 1 for details).

Each group was observed for eight consecutive hours during both days. Each ratio was presented for a total of 4 hours subdivided in two observations of 2 hours each; in one the bigger hole was presented on the frontal right side of the tank and in the other one on the frontal left side to check for any side bias. Each panel was presented once a day for each group and the sequence between groups was randomised. Before inserting a new panel, we waited until all larvae were near the two ends of the apparatus. From the video recordings, we scored the total number of passages through each hole for every panel presented for each cluster. Since both pre-test and test phases lasted two days, larvae of “7 dpf group” were 9-to-10-dpf at the moment of the cognitive measurement, those of “14 dpf group” were 16-to-17-dpf and those of “21 dpf group” were 23-to-24-dpf.

Statistical analyses

Analyses were performed in R version 3.5.2 (The R Foundation for Statistical Computing, Vienna, Austria, <http://www.r-project.org>). Each cluster of six larvae was considered as one datapoint with no distinction between the six larvae. All the data of each group are available in the supplementary file. Binomial tests were performed to compare the passages through the bigger hole in every ratio with chance level for all three ages separately and also pooled together. Post hoc power analyses were performed on the binomial tests using the ‘binom.power’ function of the ‘binom’ R package. A linear mixed-effects model (LMM, ‘lmer’ function of the ‘lme4’ R package) was performed to compare the total number of passages between the four ratios for the three larval ages. Such LMM was fitted with larvae group as random effect and with age and ratio as fixed

211 effects; both fixed effects were categorical factors. To compare the performance (passages through
212 the bigger hole) between the different ratios, between the three ages and the effect of the day, we
213 used a LMM fitted with larvae group as random effect and with age, day and ratio as fixed effects;
214 all three fixed factors were categorical factors. Subsequently, all pairwise comparisons were
215 performed with Tukey post-hoc tests. We performed another LMM, equally fitted, to compare the
216 results of the three ages with the results of the adults of the previous work (Santacà et al. 2020).
217 Partial eta-squared (ηp^2) were used as an effect size statistic for LMMs ('eta_sq' function of the
218 'sjstats' package).

219 RESULTS

220 The mean number of passages was 136 ± 63 in 16 h of recordings for the 7dpf larvae, $337 \pm$
221 144 for the 14 dpf larvae and 720 ± 287 for the 21 dpf larvae. The LMM on the total number of
222 passages revealed that the three larval ages significantly differed ($F_{(2, 27)} = 5.609$, $P < 0.01$, $\eta p^2 =$
223 0.116) while the effects of the ratio and the interaction age x ratio were not significant (ratio: $F_{(3, 81)}$
224 $= 1.665$, $P = 0.181$, $\eta p^2 = 0.055$; age x ratio: $F_{(6, 81)} = 2.144$, $P = 0.060$, $\eta p^2 = 0.131$).

225 Overall, 7 dpf larvae passed significantly more through the bigger hole in the ratios 0.60,
226 0.75, 0.86 but not in ratio 0.91; similarly, 14 dpf larvae passed significantly more through the bigger
227 hole in the ratios 0.60, 0.75, 0.86 but not in ratio 0.91 whereas 21dpf larvae passed significantly
228 more through the bigger hole in all the ratios presented (Figure 3). Table 2 shows the proportion of
229 passages through the bigger hole, 95% Confidence Interval, binomial tests and post hoc power
230 analyses for all ratios and for all ages. Pooling together the data of three ages, zebrafish larvae
231 significantly discriminate the bigger hole in all the four ratios (all P -values < 0.001).

232 A LMM revealed significant difference in performances between the four ratios ($F_{(3, 189)} =$
233 164.996, $P < 0.001$, $\eta p^2 = 0.718$). A Tukey post hoc test revealed that all the pairwise comparisons
234 between the ratios were statistically significant (all P -values < 0.05). The effects of the age and of
235 the day were not significant (age: $F_{(2, 27)} = 0.412$, $P = 0.667$, $\eta p^2 = 0.004$; day: $F_{(1, 189)} = 1.664$, $P =$
236 0.199, $\eta p^2 = 0.008$). All the interactions were not significant either (all P values > 0.333 , all $\eta p^2 <$
237 0.017).

238 Comparing the results of the larvae with the adults' result of the previous study (Santacà et
239 al. 2020), we found no significant effect of the age (LMM: $F_{(3, 32)} = 0.944$, $P = 0.431$, $\eta p^2 = 0.026$;
240 Figure 3); also the interaction age x ratio was not significant ($F_{(2, 27)} = 1.443$, $P = 0.181$, $\eta p^2 =$
241 0.108).

242

243

244 **DISCUSSION**

245 Recently we showed that adult zebrafish can be very precise in continuous quantity
246 discrimination, being able to detect also subtle size differences between two holes, such as 10%
247 difference in area (Santacà et al. 2020). In this study we investigated the ontogeny and the
248 developmental trajectory of this cognitive skill. To achieve our goal, we observed the spontaneous
249 behaviour of groups of zebrafish larvae in the presence of holes differing in size.

250 In our experiment we found that larvae of all ages prefer to pass through the larger hole. It is
251 unclear whether they used a linear measure (diameter or circumference) or the area of the hole to
252 estimate its size. In humans, the Weber function is similar for linear measures and for areas but,
253 since the surface area of a figure increases with the square of its linear size, the difference between
254 the areas of two figures is more discernible than the difference between their linear dimensions (e.g.
255 Eriksen and Hake 1955; Miller 1956; Rule 1969). Accordingly, it seems wise to consider that our
256 subjects were estimating up to a 0.91 ratio between areas rather than up to a 0.95 ratio between
257 diameters or between circumferences (see table 1).

258 There appears to be only minor differences between the three ages. Binomial tests showed
259 that 7 and 14 dpf larvae discriminated up to 0.86 ratio; only the 21 dpf larvae demonstrated to
260 discriminate the most difficult ratio (0.91) as adult zebrafish do (Santacà et al. 2020). One may be
261 tempted to conclude that the precision to make continuous quantity discrimination increases across
262 development. However, descriptive data showed quite a similar performance among the three ages
263 and LMM on the number of passages through the holes indicated that this measure increased as a
264 function of age, with 21 dpf larvae doing more than 5-fold number of passages than 7 dpf larvae.
265 This result is in accordance with previous studies showing that locomotor activity of larval
266 zebrafish steadily increases during development (e.g. Colwill and Creton 2011; Fuiman and Webb
267 1988). Therefore, the differences here reported in the binomial tests are likely to be due to the
268 different statistical power of the binomials tests rather than reflecting true differences in the
269 precision of cognitive systems underlying quantity discrimination. This was confirmed by the post

270 hoc power analyses and by the LMM on the proportion of choices for the larger hole that showed no
271 difference as a function of age even when the data of adults are included in the analyses. Our
272 experimental approach proved to be highly effective since both larvae and adults demonstrated a
273 high discrimination ability since they both discriminated between two areas that differ for less than
274 10%. To our knowledge, beside humans (Agrillo et al. 2013), such precise size quantitative
275 discrimination has been found only in sea lions trained to a size-discrimination task (Schusterman et
276 al. 1965). One subject discriminated between two geometrical shapes up to a 0.88 area ratio and the
277 other up to 0.94 ratio.

278 Interestingly, we found that the performance of all three age groups significantly decreased when
279 the ratio between the smaller and the larger hole increased, ranging from 74% (0.60 ratio) to 54%
280 (0.91 ratio). This aligns with a large body of evidence showing that the capacity to make relative
281 quantity judgments in human and non-human animals (both discrete and continuous) obeys to
282 Weber's law, which states that the ability to discriminate two stimuli could depends mainly on their
283 relative difference, rather than their absolute difference (e.g. Feigenson 2007; Ditz and Nieder
284 2016). However, our study was not designed to verify the adherence of larval zebrafish to Weber's
285 law. In fact, the size ratio and the absolute size of the smaller hole covaried so that it is difficult to
286 assess the relative influence of these two dimensions on the performance and prevent a definite
287 conclusion on a Weber's law effect. In addition, when a spontaneous preference paradigm is used, it
288 is not easy to separate the contribution of motivation and cognition (see discussion below) as they
289 are both predicted to have similar effects on performance as the ratio changed.

290 One may argue that our experiment does not allow distinguishing between the
291 discrimination and the motivation to prefer the bigger hole. Indeed, this is an important limit of all
292 cognitive studies that use spontaneous preference paradigms. For example, if an animal is observed
293 to choose the larger of two food quantities, this also implies that it can discriminate between them.
294 On the contrary, the lack of a preference could be due to either a limit in discriminability or to the
295 fact that both quantities are enough to satisfy its requirements. This factor could be a limit in our

296 study too. While it is clear that larval zebrafish are able to discriminate the bigger hole even when
297 the difference is small, our experiment cannot precisely indicate the causes of the decrease in
298 performance with increasing similarity between holes. This could be due to limits in the capacity to
299 appreciate small size differences or could be partly or entirely due to a reduced benefit of choosing
300 the larger of two holes as difference between them decreased.

301 The limitations of the spontaneous preference paradigms can be circumvented by adopting
302 an alternative procedure based on discrimination learning. In this paradigm, as motivation to obtain
303 a reward is considered independent from the stimulus to discriminate, a lack of choice should only
304 reflect the upper limit of discriminative ability. Yet, serious limitations characterize this approach
305 too, as it is time-consuming and potentially very stressful for the animal; in addition, extensive
306 training can lead to the recruitment of neurocognitive systems that are not normally involved in
307 such task (Agrillo and Bisazza 2014). In our case, a further limitation is that procedures to train
308 larval zebrafish to this task are presently unavailable.

309 Our study does not provide information about the strategy used by the larvae to solve this
310 task. There are essentially two ways that they could have used: the first one could consist in the
311 relative comparison made by visually estimating the two holes from a distance, the second one
312 could consist instead in taking an independent decision for each hole, by estimating its size using
313 vision or other sense organs (e.g., the lateral line). The former implies they possess a system for
314 relative quantity judgements, a mechanism that have been shown in the new-borns of another
315 teleost species but only for numerical judgements (Piffer et al. 2013). The latter would require that
316 they use an egocentric reference system (e.g. measuring distance between their body and the edge
317 of the hole) or in alternative comparing the size of each hole with a pre-set threshold (either innate
318 or learned). Based on what is currently known about the physiology of their visual system, it is
319 unclear whether they could have used a visual relative judgement. Adult zebrafish are known to
320 possess an excellent visual acuity (i.e. 0.56-0.58 cycles per degree; Mueller and Neuhauss 2010,
321 Tappeiner et al. 2012) but the only measurement available for larval zebrafish regards fish tested

two days after hatching, at 5dpf (Haug et al. 2010). At this stage they have poor vision, approximately one third of adults, being able to distinguish two objects only if at least 3 degrees apart. It is doubtful if this acuity would allow to perceive the small size difference proposed in our experiment (i.e. 400 μm ; see Table 1), unless they get very close to the holes, making a simultaneous estimate of the two holes impractical. However, the youngest larvae we used in our experiments were twice as old (9-10 dpf) compared to the larvae used in the cited study. It is unknown how rapidly the visual system develops but an important phase in the development of nervous system occurs around 6 dpf when larvae start to show a richer behavioural repertoire and begin to catch live preys (Fero et al. 2011). We have only indirect information about visual acuity at this stage of development. McElligott and O'Malley (2005) have studied the kinematic of prey capture in 6-8 dpf larvae feeding on paramecia. Prey capture is visually guided, and many prey tracking movements initiate when a prey is between 2.5 to 3 mm from the fish. Considering that a paramecium is 200-250 μm long, larvae of this age are certainly capable of perceiving even the smaller hole used in this study (7600 μm) from any position within the apparatus. It is unclear whether they would be able to appreciate the smallest difference between hole's diameters that was only twice the average length of a paramecium. However, if we consider that larvae likely estimated the area rather than the diameter of the holes (see discussion above) the difference in areas is well above the minimum discernible area of their prey (4.9 mm^2 and 0.02 mm^2 respectively).

In the last decade, zebrafish is rapidly gaining popularity as a model of human neuropathologies, due to a greater ease in dissecting the genetic and physiological basis of these diseases. Zebrafish models have been established for studying neurodegenerative diseases such as Alzheimer's disease, neurodevelopmental disorders such as schizophrenia, genetic or autoimmune diseases that affects nervous system such as multiple sclerosis (e.g. Kulkarni, Yellanki, Medishetti, Sriram, Saxena, Yogeeswari, 2017; Newman, Ebrahimie, Lardelli, 2014). Some of these types of neurodegenerative diseases are frequently associated with a decline in the ability to estimate quantities. The deficits involve both numeracy (e.g. Gandini et al. 2009) and the estimation of

348 continuous quantities, for example size, weight or distance of an object (e.g. Barabassy et al. 2010;
349 Brand et al. 2003). One of the advantages of studying zebrafish is the possibility to investigate
350 pathological phenomena very early, soon after birth or in the first weeks of life. A remarkable
351 example is the development of zebrafish lines with alterations of TAU protein functioning that
352 induce early neuronal disturbances and cell death, and recapitulated, in the first days of life, the key
353 pathological features of human TAU-related pathologies such as Alzheimer's disease (Paquet et al.
354 2009). Our experimental procedure has been shown in previous study on adults to have a good
355 retest reliability and to be unaffected by experience (Santacà et al. 2020). Moreover, the procedure
356 is relatively fast and records the spontaneous behaviour of fish in their home tank, avoiding
357 stressing subjects with frequent transfers from home cage to the experimental tank. Here we showed
358 that it can be administered since the first week of life. In the light of these characteristics, this test
359 appears suitable to measure longitudinal changes on cognitive abilities for example to examine the
360 effects of pharmacological treatments on neurodegenerative processes using larval zebrafish as
361 model, instead of adult zebrafish.

362

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498 **ETHICAL NOTES:** The experiments were approved by the Ethical Committee of the Università
499 di Padova (protocol n. 61/2018) and adhered to the current legislation of the country in which they
500 were performed (Italy, Decreto Legislativo 4 Marzo 2014, n. 26).

501 **CONFLICT OF INTEREST STATEMENT:** We declare no conflict of interests.

502 **DATA AVAILABILITY STATEMENT:** The data that support the findings of this study are
503 available from the corresponding author upon request.

504

505 **TABLE**

506 **Table 1. Hole diameters and size ratios used in this study.**

Diameter of larger hole	Diameter of smaller hole	Ratio between the diameters	Ratio between the areas
0.8 cm	0.62 cm	0.78	0.60
0.8 cm	0.69 cm	0.86	0.75
0.8 cm	0.74 cm	0.93	0.86
0.8 cm	0.76 cm	0.95	0.91

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508 **Table 2. Performance of the three ages.** Mean proportion of choices for the bigger hole, 95%
509 Confidence Interval and Binomial Tests for all ratios and for the three ages.

510					
Larval age	Ratio	Proportion of choices for the bigger hole (mean \pm SD)	95% Confidence Interval	Binomial Tests	Post hoc power analyses
7 dpf	0.60	0.766 \pm 0.034	0.742, 0.790	$P < 0.05$	$1 - \beta = 1.000$
	0.75	0.636 \pm 0.072	0.585, 0.687	$P < 0.05$	$1 - \beta = 1.000$
	0.86	0.594 \pm 0.071	0.543, 0.644	$P < 0.05$	$1 - \beta = 0.870$
	0.91	0.549 \pm 0.071	0.507, 0.582	$P = 0.073$	$1 - \beta = 0.280$
14 dpf	0.60	0.736 \pm 0.066	0.689, 0.783	$P < 0.001$	$1 - \beta = 1.000$
	0.75	0.649 \pm 0.069	0.600, 0.699	$P < 0.001$	$1 - \beta = 1.000$
	0.86	0.568 \pm 0.064	0.522, 0.614	$P < 0.001$	$1 - \beta = 0.901$
	0.91	0.519 \pm 0.074	0.466, 0.572	$P = 0.124$	$1 - \beta = 0.152$
21dpf	0.60	0.716 \pm 0.062	0.671, 0.760	$P < 0.001$	$1 - \beta = 1.000$
	0.75	0.642 \pm 0.075	0.589, 0.696	$P < 0.001$	$1 - \beta = 1.000$
	0.86	0.594 \pm 0.026	0.575, 0.612	$P < 0.001$	$1 - \beta = 1.000$
	0.91	0.546 \pm 0.046	0.513, 0.578	$P < 0.001$	$1 - \beta = 0.980$

514 **FIGURES**

515 **Figure 1. Experimental apparatus.** a) Lateral and b) aerial view of the experimental context. The
516 apparatus was composed of a movable test panel in the middle of the corridor that divided the
517 apparatus in a frontal and a posterior compartment.

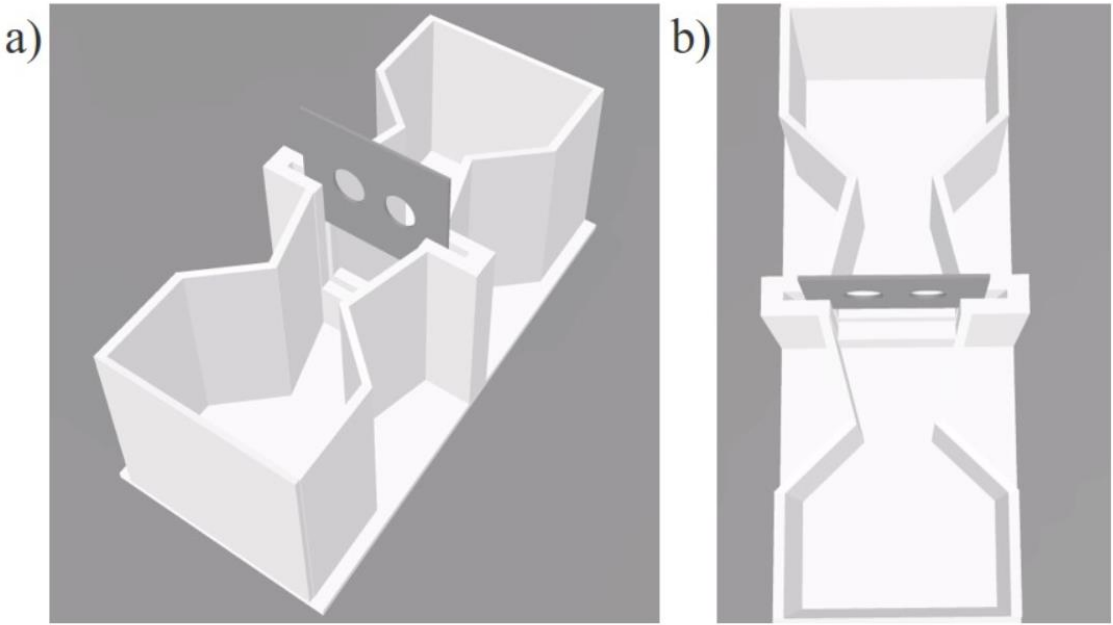
518 **Figure 2. Experimental panels.** Subjects were presented with panels showing pairs of holes with
519 four different ratios between the areas within each pair: ratio 0.60 (a), ratio 0.75 (b), ratio 0.86 (c)
520 and ratio 0.91 (d).

521 **Figure 3. Comparison of the results of the three larval ages and the adults of the previous**
522 **work (Santacà et al. 2020).** The Y-axis refers to the proportion of choices for the bigger hole in the
523 four ratios tested (ratio 0.60, ratio 0.75, ratio 0.86, ratio 0.91). Bars represent the standard error.

524

525 **Figure 1.**

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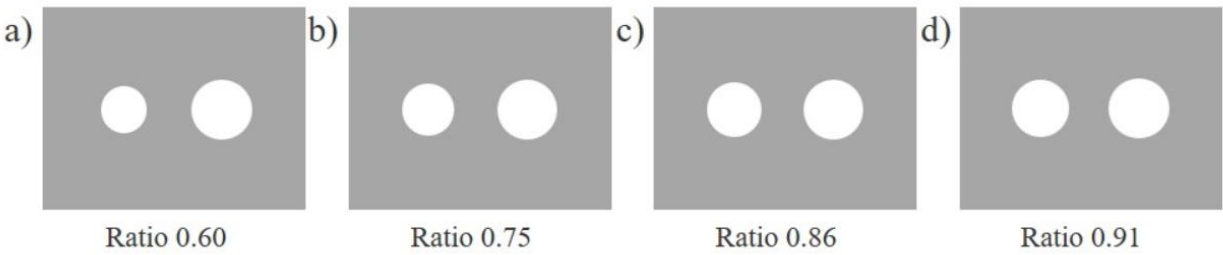
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535 **Figure 2.**

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549 **Figure 3.**

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